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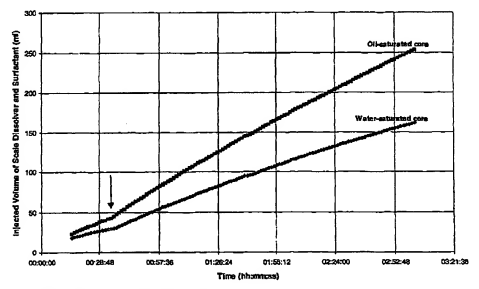
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[Continued on next page]

(54) Title: SCALE DISSOLVER FLUID



(57) Abstract: A scale dissolver fluid for dissolving scale in a subterranean hydrocarbon-bearing formation comprises an effective amount of a scale dissolver formulation and an effective amount of a surfactant for controlling the viscosity of the fluid. In use, formation hydrocarbons act on the surfactant to reduce the viscosity of the fluid so that the fluid selectively invades a hydrocarbon-bearing zone of the formation.

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#### SCALE DISSOLVER FLUID

#### Field of the Invention

The present invention relates to a scale dissolver

fluid for dissolving scale in hydrocarbon wells, and to a
method of dissolving scale in hydrocarbon wells.

#### Background of the Invention

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The recovery of hydrocarbons, such as oil and gas,

from a subterranean well formation can be impeded by
scales obstructing the flow of hydrocarbons from
hydrocarbon-bearing zones of the formation. Typical
scales are barite (e.g. BaSO<sub>4</sub>) or calcite (e.g. CaCO<sub>3</sub>) and
it is common practice to treat these by bull-heading an
aqueous-based scale dissolver fluid through a well bore
and into the formation.

For example, one conventional scale dissolver for barite scale consists of a concentrated solution of potassium carbonate, potassium hydroxide and the potassium salt of ethylenediaminetetraacetic acid(EDTA), the corrosive and chelating nature of the solution being effective in removing scale. Carbonate scales may be dissolved using simple mineral acids, such as HCl.

However, hydrocarbon-producing wells often contain zones that are watered-out, producing only, or very largely, water. If the scale dissolver enters these zones, scale may also be removed therefrom. This can lead to an undesirable increase in the water cut of the fluid produced by the well.

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In related but different fields of hydrocarbon recovery (notably the field of hydraulic fracturing, as described for example in EP-A-0835983), significant use is made of viscoelastic fluids. These fluids are typically based on aqueous solutions of surfactants, such as erucyl bis(2-hydroxyethyl) methyl ammonium chloride or potassium oleate, which can form worm-like micelles when mixed with brines, e.g. KCl brine. The structure of the micelles contributes significantly to the viscoelasticity of the fluid, and viscoelasticity is rapidly lost when the fluid contacts hydrocarbons which cause the micelles to change structure or disband.

#### Summary of the Invention

In accordance with the present invention a scale dissolver fluid is provided for dissolving scale in a hydrocarbon well, the fluid including means for controlling the viscosity of the fluid.

A first aspect of the present invention provides a scale dissolver fluid for dissolving scale in a subterranean hydrocarbon-bearing formation, the fluid comprising an effective amount of a scale dissolver formulation and an effective amount of a surfactant for controlling the viscosity of the fluid, whereby in use formation hydrocarbons act on the surfactant to reduce the viscosity of the fluid so that the fluid selectively invades a hydrocarbon-bearing zone of the formation.

In use the fluid is injected into the subterranean formation in a relatively viscous state. If the injected fluid contacts a watered-out zone of the formation the

viscous nature of the fluid remains essentially unaltered and, to a significant extent, the fluid is prevented from entering the watered-out zone, i.e. the fluid locally has limited injectivity. Conversely, if the fluid contacts a hydrocarbon-bearing zone of the formation the viscosity is locally significantly reduced and the fluid is able to penetrate the hydrocarbon-bearing zone.

Therefore, the difference in viscosity of the fluid when in contact with hydrocarbons and water

10 advantageously allows a selective placement of the scale treatment, and as a result scale may be preferentially removed from hydrocarbon-bearing zones. This can lead to a stimulation of hydrocarbon production without a substantial increase in the water cut of produced fluids.

by reversibly producing viscoelasticity in the fluid.

That is, the fluid is viscoelastic in nature when injected and this property is maintained in aqueous environments (e.g. watered-out zones). However,

formation hydrocarbons act on the surfactant to destroy or reduce the viscoelasticity, allowing the fluid to penetrate hydrocarbon-bearing zones.

By "viscoelastic", we mean that the elastic (or storage) modulus G' of the fluid is greater than the loss modulus G" as measured using an oscillatory shear rheometer (such as a Bohlin CVO 50) at a frequency of 1 Hz. The measurement of these moduli is described in An Introduction to Rheology, by H.A. Barnes, J.F. Hutton, and K. Walters, Elsevier, Amsterdam (1997).

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More preferably the scale dissolver formulation activates the production of viscoelasticity by the surfactant. In this way it may not be necessary to add additional agents, such as KCl brine, to activate the production of viscoelasticity. However, the use of such additional agents is not excluded by the present invention. The scale dissolver formulation may comprise any acid or alkaline solution that dissolves minerals and other well bore deposits (including organic deposits).

Desirably the scale dissolver formulation comprises an

aqueous solution of at least one of an alkali metal carbonate, alkali metal hydroxide, EDTA and the alkali metal salt of EDTA. The alkali metal may be potassium. Alternatively the scale dissolver formulation may comprise a mineral acid, such as HC1.

The surfactant of the scale dissolver fluid may be any anionic or cationic surfactant that forms a viscoelastic gel in aqueous media and whose viscoelasticity is reduced or destroyed on contact with hydrocarbons. For example the surfactant may comprise Nerucyl-N,N-bis(2-hydroxyethyl)-N-methyl ammonium chloride, a salt of oleic acid (e.g. an alkali metal salt such as potassium oleate), or a salt of an oligomer of oleic acid (e.g. an alkali metal salt, such as a potassium salt). The oleic acid salt and oleic acid oligomer salt may be formed in situ from the corresponding acid precursors. By an "oligomer of oleic acid" we mean an unhydrogenated, a fully hydrogenated or a partially hydrogenated oligomer of oleic acid.

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A second aspect of the invention provides a scale dissolver fluid for dissolving scale in a subterranean formation, the fluid comprising an aqueous solution of at least one scale dissolving component and at least one surfactant, and having substantially Newtonian viscous behaviour at least in the shear rate range 0.1-100 (preferably 0.1-1000) s<sup>-1</sup> and a viscosity in the range 20 to 1000 (preferably 100 to 1000) centipoise at 60°C, the viscosity falling to a value in the range 1 to 200 (preferably 1 to 50) centipoise on contact with a hydrocarbon fluid, such as heptane, mineral spirits or crude oil.

Depending on the type of scales, fluids of different pH might be suitable. For instance, for barium and strontium sulphate, fluids having a pH greater than 12 are usually required. Anionic surfactants (for examplesalts of oleic acid or of an oligomer of oleic acid e.g. an alkali metal salt, such as a potassium salt) compatible with such high pH, are therefore preferred. For calcium sulphate, a pH ranging from 7 to 10 is suitable, a preferred option is a cationic surfactant. This is also the case with calcium carbonate, where the pH has preferably to be optimised close to 5.

A third aspect of the present invention provides a

25 method of dissolving scale in a subterranean formation
with at least one hydrocarbon-bearing zone, the method
including pumping the scale dissolver fluid of the first
or second aspect of the invention through a well bore and
into the subterranean formation, the viscosity of the

30 scale dissolver fluid being reduced by formation

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hydrocarbons so that the fluid selectively invades the hydrocarbon-bearing zone of the well to dissolve scale in the hydrocarbon-bearing zone

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A fourth aspect of the present invention provides a method of injecting a scale dissolver fluid into a subterranean formation with at least one hydrocarbonbearing zone, the method including the step of pumping the scale dissolver fluid of the first or second aspect of the invention through a well bore and into the subterranean formation.

In both the third and fourth aspects of the invention, preferably the scale dissolver fluid is bullheaded through the well bore.

#### 15 Brief Description of the Drawings

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Specific embodiments of the present invention will now be described with reference to the following drawings in which:

Fig. 1 shows a graph of viscosity against shear 20 rate at various temperatures for a scale dissolver fluid containing N-erucyl-N,N-bis(2-hydroxyethyl)-N-methyl ammonium chloride;

Fig. 2 shows a graph of viscosity against shear rate at various temperatures for a scale dissolver fluid 25 containing potassium oleate;

Figs 3a-e show various oleic acid dimers;

Fig. 4 shows a graph comparing the rheology of two scale dissolver fluids comprising oleic acid oligomers at 60°C;

- Fig. 5 shows a graph comparing the injectivities into oil and water-saturated cores of a scale dissolver fluid;
- Fig. 6 shows schematically the steps involved in deploying a scale dissolver fluid of the present invention;
  - Fig. 7 shows a plot of the amount dissolved (g/1) versus the incubation time;
- Fig. 8 shows the fluid expelled from each core

  10 expressed as a fraction of total fluid expelled from both
  cores:
  - Fig. 9 shows the fractional diversion in the low permeability cores; and
- Fig. 10 shows the fluid expelled from each core

  15 expressed as a fraction of total fluid expelled form both
  cores.

#### Detailed Description

The scale dissolver fluid of the present invention

20 has an enhanced rheological performance which allows it
to dissolve scales preferentially in hydrocarbon-bearing
matrices of subterranean formations. To a significant
extent this performance is due to the ability of the
fluid to vary its viscosity depending on whether it is in

25 contact with water or hydrocarbons. In contrast,
conventional scale dissolver fluids remove scale deposits
indiscriminately from hydrocarbon and water-bearing zones
alike.

Scale dissolver fluids of the present invention may 30 contain viscoelastic surfactants for forming viscoelastic

gels. If the fluid is considered as a combination of a conventional scale dissolver fluid and such a surfactant, the viscosity of the gel can be reduced to substantially that of the conventional fluid when the gel comes into contact with hydrocarbons, making the scale dissolver formulation of the fluid readily injectable into hydrocarbon-bearing matrices. However, when the gel contacts water it remains highly viscous (and therefore not easily injectable), any reduction in viscosity being essentially due to dilution. Effectively the highly viscous gel acts as a diverting agent and allows a high proportion of the scale dissolver formulation to be placed in hydrocarbon zones.

#### 15 Example 1

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13g of EDTA, 11.25g of potassium hydroxide and 2.25g of potassium carbonate were dissolved in 70.5g of water. 3g of N-erucyl-N,N-bis(2-hydroxyethyl)-N-methyl ammonium chloride was then added and the mixture stirred until a homogeneous gel was formed.

#### Example 2

6.5g of EDTA, 5.625g of potassium hydroxide and 1.125g of potassium carbonate were dissolved in 83.75g of water. 3g of potassium oleate was then added and the mixture stirred until a homogeneous gel was formed.

In Examples 1 and 2 the potassium hydroxide and the potassium carbonate activated the production of viscoelasticity by the N-erucyl-N,N-bis(2-hydroxyethyl)-

N-methyl ammonium chloride and potassium oleate respectively.

A controlled stress rheometer (Bohlin model type CVO-50) was used to measure the rheological properties of the systems of Examples 1 and 2. Using a concentric cylinders (Couette) geometry (inner radius of the outer cylinder,  $R_i = 1.375$ cm, outer radius of the inner cylinder, Ro = 1.25cm, and inner cylinder length = 3.78cm), corresponding to the geometry of German DIN 10 standard 53019, the viscosity of the sample was measured at several applied shear stresses within a specified range. The typical range of shear stress was 0.5-40 Pa corresponding to a shear rate range of 0.005 to 1000s<sup>-1</sup>. Measurements were made at increasing and then decreasing shear rate. Typically, the complete set of measurements 15 consisted of 40 viscosity measurements, each taken after a delay time of 10 seconds at constant shear stress and shear rate.

For the particular geometry of the rheometer, the 20 shear rate was calculated as:

$$\dot{\gamma} = \frac{RPM.2\pi}{60} \frac{2.R_i^2 R_0^2}{\left(\frac{R_i + R_0}{2}\right)^2 \left(R_0^2 - R_i^2\right)},$$

where RPM is the relative rotational speed (in revolutions per minute) of the cylinders. The viscosity was then obtained for each measurement by dividing the measured stress by the calculated shear rate.

Figs. 1 and 2 respectively illustrate the rheology of the systems of Examples 1 and 2 measured in this way at various temperatures in the range 25-80°C. Varying the amount of surfactant or changing the types of inorganic ions can vary the rheology of each gel so that the gel can be optimised for specific applications.

#### Example 3

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The oligomerisation of oleic acid generally leads to the production of complex mixtures of dimeric and 10 trimeric products. Commercially available mixtures, such as the Empol™ series of dimers and trimers from Henkel Corporations Chemical Group (4900 Este Avenue-Bldg 53, Cincinnati, Ohio 45232, USA) are suitable for putting the present invention into operation. Alternative suppliers 15 of suitable mixtures are e.g. Union Camp (Vigo Lane, Chester-le-Street. Co. Durham DH3 2RB, UK) and Expo Chemical Company Inc. (12602 Manorwood, Cypress (Houston), Texas 77429, USA). Figs. 3a-e show typical chemical structures of dimeric components of these 20 mixtures. Clearly the components have different degrees of hydrogenation.

EDTA (13g), potassium hydroxide (11.25g) and potassium carbonate (2.25g) were dissolved in water (70.5g). Empol<sup>TM</sup> 1016 dimer acid (3g) was then added and the mixture stirred until it became a homogeneous gel.

#### Example 4

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EDTA (8.66g), potassium hydroxide (7.5g) and 30 potassium carbonate (1.5g) were dissolved in water (79g).

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Empol<sup>TM</sup> 1043 trimer acid (3g) was then added and the mixture stirred until it became a homogeneous gel.

The viscosities of the gels of Examples 3 and 4 were measured (using the procedure described above for Examples 1 and 2) at 60°C over a range of shear rates. The results of these measurements are shown in Fig. 4. Both gels exhibited Newtonian rheology over a surprisingly wide range of shear rates. Advantageously, therefore, the injectivity of the gels into subterranean matrices should not be affected by changes in shear rate which may occur during the placement process. However, the viscosity of a scale dissolver fluid containing a mixture of such oligomers can be controlled by adjusting the amount and type of oligomer in the mixture.

A 150 cP gel based on the formulation of Example 3 was injected into an oil-saturated core and a water-saturated core by forcing the gel down a supply line which branched into two parallel lines leading to the two cores. Both cores were of Bentheimer sandstone and had equal total pore volumes. By measuring the relative amounts of gel entering the two cores at a given supply pressure or for a given volume of supplied gel, the relative injectivities of the gel through the two cores was determined.

25 Injection profiles of the gel into the two cores with the fluid and cores maintained at a temperature of 60°C are shown in Fig. 5. The permeability of the water-saturated core was 1.6 darcies while that of the oil-saturated core was 1.4 darcies; both cores had a porosity of 22%. The profiles demonstrate that the volume of gel

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entering the oil-saturated core is approximately 50% greater than that entering the water-saturated core. The preference of the gel to enter the oil-saturated core is maintained even after a large number of pore volumes was passed through the two cores. The viscosity of the effluent from the oil-saturated core was significantly lower than that of the injected gel throughout the duration of the experiment and demonstrated that the surfactant gel was continually mix with oil. In contrast, the viscosity of the effluent from the water-saturated core was similar to that of the injected gel. Higher viscosity fluids enhance this contrast and fluids can be developed that only enter oil-bearing zones, the viscosity being too high for injection into the water-bearing zones.

Fig. 6 shows schematically the steps involved in the deployment of a scale dissolver fluid of the present invention.

While the invention has been described in conjunction

with the exemplary embodiments described above, many
equivalent modifications and variations will be apparent
to those skilled in the art when given this disclosure.

Accordingly, the exemplary embodiments of the invention
set forth above are considered to be illustrative and not
limiting.

#### Example 5

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A viscoelastic fluid, composed of 8.1g ethylenediaminetetraacetic acid, 7.22g potassium hydroxide, 1.4g potassium carbonate, 2g oleic acid and 81.3g of water, was constructed. To this fluid was added

dissolver used as a benchmark.

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5ml of diesel oil in order to reduce viscosity; this process is known as breaking the fluid. To 10ml of this reduced viscosity fluid was added to an exactly known mass of crystalline barium sulphate M1 (mass approximately 1g) with an approximate mean particle size of 100 m. mixture was incubated in a plastic bottle at 50°C for 1 hour, after which the solid barium sulphate was removed by decantation, washed with distilled water, dried at 70°C and the new mass  $M_2$  determined. The difference between  $M_1$ and  $M_2$  was used to compute the amount dissolved in g/1 at the specified time. This experiment was repeated for incubation times of 1hr, 2hr, 4hr, 8hr, 16hr, 20hr and Fig. 7 shows a plot of the amount dissolved (g/1) versus the incubation time. Data are compared to an identical suite of measurements collected using 2butoxyethanol as an alternative viscosity reducer to diesel, and also to data collected using the dissolver formulation made without oleic acid. This later formulation is simply a surfactant-free chelating

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Inspection of Fig. 7 demonstrates that a dissolver fluid, having once been viscous for the purpose of diversion but now broken, can dissolve barium sulphate with an efficiency close to that of a benchmark dissolver.

#### Example 6

An experimetal setup was constructed in order to inject a viscoelastic scale dissolver fluid through two fluid saturated cores simultaneously. Both cores had

permeabilities of approximately 50mD, but one core was saturated to 80% with hydrocarbon and 20% with water, the other was saturated to 80% with water and 20% with diesel The viscoelastic scale dissolver was composed of 9g oil. ethylenediaminetetraacetic acid, 6.8g potassium hydroxide, 2g oleic acid and 100g of water. This fluid was injected simultaneously through both cores, through a supply line that branched into two parallel lines leading into each core. Tests were conducted at 50°C using 12 inch cores with a differential pressure of 100psi. 10 Immediately upon onset of injection the mass of fluid expelled from each core was determined, by weighing the expelled fluid, continuously as a function of time over a period of 25 minutes. Fig. 8 shows the fluid expelled from each core expressed as a fraction of total fluid 15 expelled from both cores.

This shows that over 80% of the injected fluid can be diverted through a predominantly oil-bearing core, despite the fact that this had approximately 20% residual water and the water bearing core had approximately 20% diesel oil. The presence of hydrocarbon in the water core is expected to break the fluid to some extent, and the presence of water in the oil core is expected to limit breaking, both factors will act against diversion. The high level of diversion, even in these cases, indicates that diversion can be maintained with a suitable fluid formulation.

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This experiment was repeated using a viscoelastic scale dissolver composed of 9g ethylenediaminetetraacetic acid, 6.8g potassium hydroxide, 2g oleic acid, 1g of a

sugar surfactant (an alkyl polyglycoside of CAS number 68515-73-1) and 100g of water. The core permeabilities were approximately 700mD.

Fig. 9 shows the fractional diversion in the low permeability cores. With this formulation, a reduced diversion was observed but the data indicates that alternative surfactants may be compatible with scale dissolver formulations.

#### 10 Example 7

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An experimental setup was constructed in order to inject a viscoelastic scale dissolver fluid independently through two cores having a permeability contrast of over one order of magnitude. The low permeability core (50mD) was saturated to 100% with diesel oil and the high permeability core (700mD) was saturated to 100% with water. The viscoelastic scale dissolver was composed of 9g ethylenediaminetetraacetic acid, 6.8g potassium hydroxide, 2g oleic acid and 100g of water. This fluid was injected independently through each core using a single supply line into the respective core. The independent use of cores differentiates this test from that in Example 7, which involved simultaneous core flooding. Tests were conducted at 50°C using 12inch cores with a differential pressure of 100psi. Immediately upon onset of injection the mass of fluid expelled from each core was determined, by weighing the expelled fluid, continuously as a function of time over a period of 25 minutes.

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Fig. 10 shows the fluid expelled from each core expressed as a fraction of total fluid expelled form both cores. This shows that over 65% of the injected fluid can be diverted through an oil-bearing core despite the fact that the alternative flow path had a massively preferential permeability. The high level of diversion, even in these cases, implies that diversion can be achieved with high permeability contrasts using a suitable fluid formulation.

Various changes to the described embodiments may be made without departing from the spirit and scope of the invention

#### Claims

- 1. A scale dissolver fluid for dissolving scale in a subterranean hydrocarbon-bearing formation, the fluid comprising an effective amount of a scale dissolver formulation and an effective amount of a surfactant for controlling the viscosity of the fluid, whereby in use formation hydrocarbons act on the surfactant to reduce the viscosity of the fluid so that the fluid selectively invades a hydrocarbon-bearing zone of the formation.
  - 2. A scale dissolver fluid according to claim 1, wherein the surfactant controls the viscosity of the fluid by reversibly producing viscoelasticity in the fluid.
  - 3. A scale dissolver fluid according to claim 2, wherein the scale dissolver formulation activates the production of viscoelasticity by the surfactant.

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- 4. A scale dissolver fluid according to any one of claims 1 to 3, wherein the surfactant comprises an oleic acid salt or a salt of an oligomer of oleic acid.
- 5. A scale dissolver fluid according to any one of claims 1 to 4, wherein the scale dissolver formulation comprises an aqueous solution of at least one of potassium carbonate, potassium hydroxide, ethylenediaminetetraacetic acid (EDTA) and the potassium salt of EDTA.

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6. A scale dissolver fluid according to any one of claims 1 to 3, wherein the surfactant comprises N-erucyl-N,N-bis(2-hydroxyethyl)-N-methyl ammonium chloride.

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- 7. A scale dissolver fluid according to any one of claims 1 to 3 and 6, wherein the scale dissolver formulation comprises an aqueous solution of HCl.
- 8. A scale dissolver fluid for dissolving scale in a subterranean formation, the fluid comprising an aqueous solution of at least one scale dissolving component and at least one surfactant, and having substantially Newtonian viscous behaviour at least in the shear rate range 0.1-100 s<sup>-1</sup> and a viscosity in the range 20 to 1000 centipoise at 60°C, the viscosity falling to a value in the range 1 to 200 centipoise on contact with a hydrocarbon fluid.
- 9. A method of dissolving scale in a subterranean formation with at least one hydrocarbon-bearing zone, the method including pumping the scale dissolver fluid of any one of claims 1 to 8 through a well bore and into the subterranean formation, the viscosity of the scale dissolver fluid being reduced by formation hydrocarbons so that the fluid selectively invades the hydrocarbon-bearing zone of the well to dissolve scale in the hydrocarbon-bearing zone.

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- 10. A method of dissolving scale in a subterranean formation according to claim 9, wherein the scale dissolver fluid is bull-headed through the well bore.
- 11. A method of injecting a scale dissolver fluid into a subterranean formation with at least one hydrocarbon-bearing zone, the method including the step of pumping the scale dissolver fluid of any one of claims 1 to 8 through a well bore and into the subterranean formation.
  - 12. A method of injecting a scale dissolver fluid according to claim 11, wherein the scale dissolver fluid is bull-headed through the well bore.

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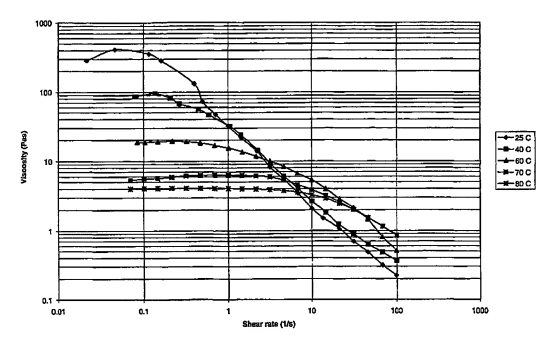


Fig. 1

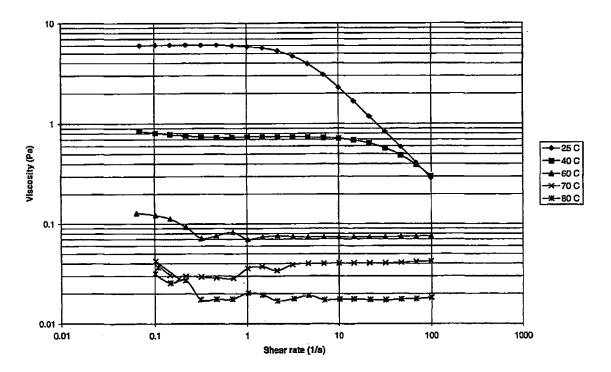


Fig. 2

Fig. 3

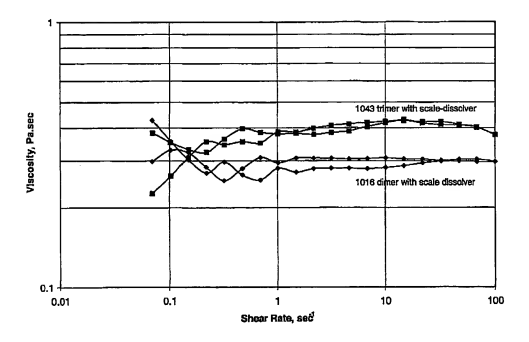


Fig. 4

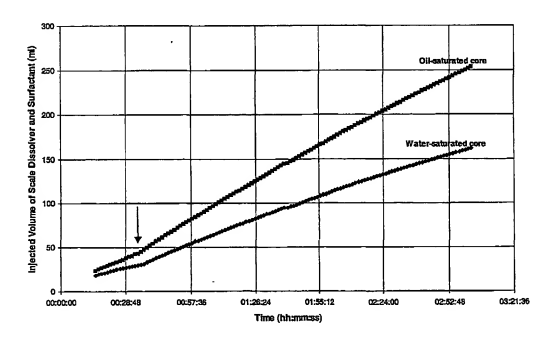


Fig. 5

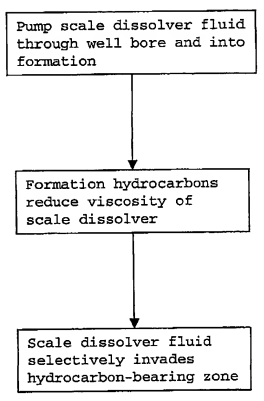


Fig. 6

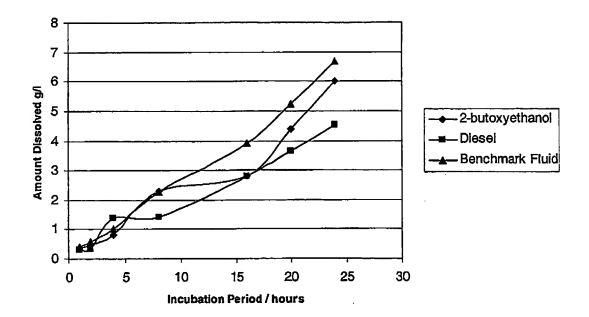


Fig. 7

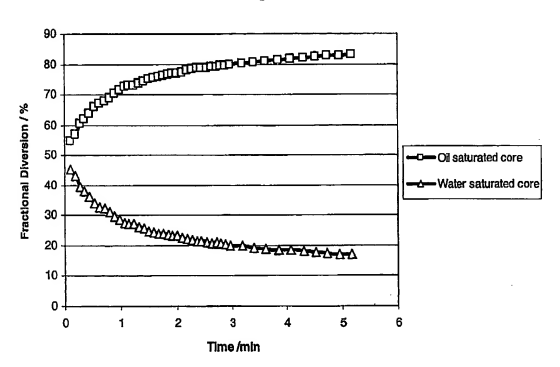


Fig. 8

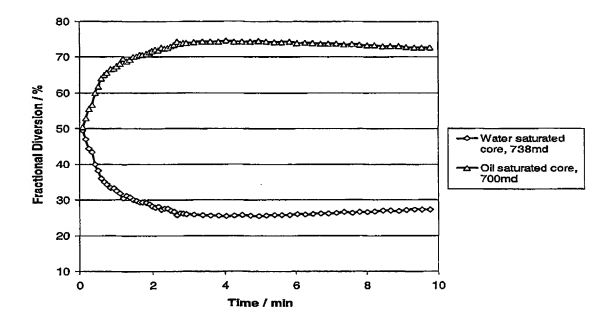


Fig. 9

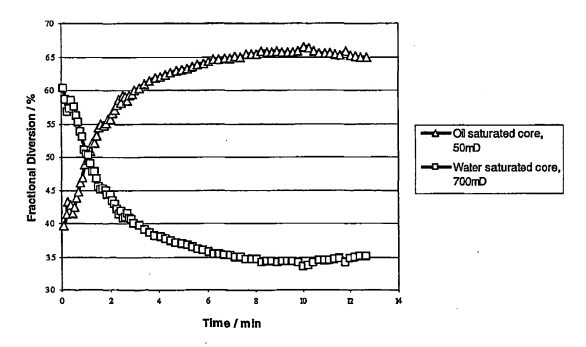


Fig. 10

#### INTERNATIONAL SEARCH REPORT

Intel nal Application No PCT/GB 01/03294

# A. CLASSIFICATION OF SUBJECT MATTER IPC 7 E21B37/06

According to International Patent Classification (IPC) or to both national classification and IPC

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 E21B B01F

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the International search (name of data base and, where practical, search terms used)

| C. DOCUMENTS CONSIDERED TO BE RELEVANT |   |                       |  |  |  |  |  |
|--|---|-----------------------|--|--|--|--|--|
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| Special categories of cited documents:  A* document defining the general state of the art which is not considered to be of particular relevance  E* earlier document but published on or after the international filing date  L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)  O* document referring to an oral disclosure, use, exhibition or other means  P* document published prior to the international filing date but later than the priority date claimed | "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention  "X" document of particular relevance; the ctaimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone  "Y" document of particular relevance; the ctaimed invention cannot be considered to involve an inventive step when the document is considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.  "&" document member of the same patent family |
| Date of the actual completion of the international search  7 November 2001   | Date of mailing of the international search report  19/11/2001   |
| Name and mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2  NL – 2280 HV Rijswijk  Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,  Fax: (+31-70) 940-3016   | Authorized officer  Boulon, A  |

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